REMARKS/ARGUMENTS

In view of the remarks and arguments below, Applicant believes the pending application is in condition for allowance.

I. Status of the Claims

Claims 1 and 3-27 were previously pending.

Claims 2, 28, and 29 were previously canceled without prejudice to or disclaimer of the subject matter contained therein.

Claims 7, 8, 10, 13, 21, 22, and 26 are canceled in this Amendment without prejudice to or disclaimer of the subject matter contained therein.

Claims 1, 14, and 16 are amended as follows.

Claims 1 and 16 are amended to change the range of the variable m recited from "2 to 100" to "10 to 100" (i.e., "m represents an integer from [[2]]10 to 100"). Support for the amendments can be found, for example, in canceled claim 8 for claim 1, and in canceled claim 22 for claim 16. No new matter is introduced by the amendments.

Claims 1 and 16 are further amended to recite that "the copolymer exhibits a microphase separated structure." Support for the amendments can be found, for example, in canceled claim 13 for claim 1, and in canceled claim 26 for claim 16. No new matter is introduced by the amendments.

Claim 14 is amended to place the Markush language recited in proper format (i.e., "selected from [[al]the group consisting of"). No new matter is introduced by the amendment.

Upon entry of this Amendment, claims 1, 3-6, 9, 11, 12, 14-20, 23-25, and 27 are pending and at issue

II.

Withdrawal of Final Office Action Dated April 9, 2008

Applicant appreciatively thanks Examiner Bernshtevn for withdrawing the Final Office Action dated April 9, 2008, and for issuing in its place the currently outstanding Non-Final Office Action dated May 13, 2008.1

III. Claim Rejections under 35 U.S.C. § 103(a)

Claims 1 and 3-27 are rejected under 35 U.S.C. § 103(a) as unpatentable over a journal article authored by Khan et al.2 ("Khan") in view of United States Patent No. 5,196,484 to Giles et al. ("Giles"). The Examiner states that Khan in combination with Giles renders the claims obvious.

Claims 7, 8, 10, 13, 21, 22, and 26 have been canceled, rendering the rejections of these claims moot.

For the pending claims 1, 3-6, 9, 11, 12, 14-20, 23-25, and 27, Applicant respectfully traverses the rejections at least for the reasons that: (A) the present invention achieves unexpected, superior results that cannot be predicted from Khan and Giles, either alone or in combination; and (B) regardless of the results achieved, one of ordinary skill in the art would not be motivated to combine Khan and Giles.

A. Unexpected, Superior Results Achieved by the Present Invention

The Examiner's rationale for establishing a prima facie case of obviousness relies on the alleged close structural similarity between the copolymers of the present invention and those suggested by the combination of Khan and Giles. However, "[a] prima facie case of obviousness based on structural similarity is rebuttable by proof that the claimed compounds possess

Docket No : 20241/0202402-US0

¹ Non-Final Office Action dated May 13, 2008, page 2, lines 13-14.

² Ishrat M. Khan, Daryle Fish, Yadollah Delayiz, and Johannes Smid, "ABA Triblock Comb Copolymers with Oligo(Oxyethylene) Side Chains as Matrix for Ion Transport," Makromolekular Chemie, 190, 1069-78 (1989).

^{3 &}quot;When chemical compounds have very close structural similarities and similar utilities, without more a prima facie case [of obviousness] may be made." Non-Final Office Action dated May 13, 2008, page 3, lines 6-8 (quoting In re Wilder, 563 F.2d 457, 461 (CCPA 1977)).

unexpectedly advantageous or superior properties." Applicant respectfully submits that such proof was previously presented.

Specifically, Applicant previously presented evidence that the claimed arrangement of copolymer block chains was nonobvious based on their unexpectedly superior conductive properties compared to the prior art. The room temperature conductivities of copolymer-LiClO4 electrolyte complexes disclosed by Khan range from 0.7 to $1.2 \times 10^{-6} \, \Omega^{-1} \cdot \mathrm{cm}^{-1}$ (Table 4). In contrast, experimental data show that the presently claimed copolymer-LiClO4 electrolyte complexes exhibit conductivities that are greater by more than an order of magnitude — from 5×10^{-5} to $3.8 \times 10^{-4} \, \Omega^{-1} \cdot \mathrm{cm}^{-1}$ (Examples 1-4 of the Specification).

In response, the Examiner stated that "it is worth to mention that Khan discloses that the ion conduction can reach values of $10^{-4} \, \Omega^{-1} \cdot \mathrm{cm}^{-1}$ at $70^{\circ}\mathrm{C}$, depending on salt and styrene content (abstract)." However, Khan states that "[a]ddition of dimethyltetraethyleneglycol (2.5.8.11,14-pentaoxapentadecane) enhances the ion conduction which can reach values of $10^{-4} \, \Omega^{-1} \cdot \mathrm{cm}^{-1}$ at $70^{\circ}\mathrm{C}$, depending on salt and styrene content" (Abstract; emphases added).

Adding DMTEG, ⁷ a plasticizer, to a polymer electrolyte and increasing temperature increases the conductivity of the electrolyte (somewhat at the expense of the mechanical properties of the polymer due to the plasticizer). ⁸ In contrast, Applicant measured the conductivities of the presently claimed solid polymer electrolytes without DMTEG and at room temperature.

Accordingly, conductivity measured (1) after the addition of dimethyltetraethylene glycol (DMTEG) and (2) at a much higher temperature does not relate to the conductivity measured by Applicant, and fails to show that the results of the present invention were ordinary or expected. The

⁴ MPEP § 2144.09, VII, first sentence.

⁵ Amendment filed on May 30, 2007, pages 10-11.

 $^{^{\}rm 6}$ Final Office Action dated August 23, 2007, page 6, lines 4-6.

⁷ The acronym "DMTEG" is a shorthand notation for dimethyltetraethyleneglycol (2,5,8,11,14-pentaoxapentadecane). Khan, p. 1077, line 14.

⁸ Khan, page 1077, second paragraph and Table 3.

Application No. 10/523,085 Amendment dated September 12, 2008 Reply to Non-Final Office Action of May 13, 2008

nature of Applicant's results should only be compared to the data from Khan at room temperature and in the absence of DMTEG — from 0.7 to $1.2 \times 10^{-6} \, \Omega^{-1} \cdot \text{cm}^{-1}$ (Table 4).

Moreover, neither Khan nor Giles suggests that rearranging the block chains of Khan would improve conductivity, let alone to the degree discovered in the present invention. In fact, as previously argued, ⁹ Khan teaches away from this rearrangement. Khan states that increasing the amount of the polystyrene block will lower the conductivity. Accordingly, the conductivities of the presently claimed solid polymer electrolytes were not only superior to the those of the prior art but also unexpected. KSR Int'l Co. v. Teleflex Inc., 127 S. Ct. 1727, 1740 (2007) ("when the prior art teaches away from combining certain known elements, discovery of successful means of combining them is more likely to be nonobvious").

At least for the reasons stated above, Applicant respectfully submits that Khan in view of Giles does not render any of claims 1, 3-6, 9, 11, 12, 14-20, 23-25, and 27 obvious, and respectfully requests that the rejections of these claims be withdrawn.

B. Lack of Motivation to Combine Khan and Giles

Further, according to the Examiner, the only difference between Khan and the present invention is the sequence of the block chains A, B, and C. ¹⁰ The Examiner's position is that it would have been obvious to one having ordinary skill in the art to place the A block in the middle of the Khan ABA block (i.e., to form a BAB block copolymer) as taught by Giles in order to reduce ambient temperature crystallization. ¹¹

However, the motivation cited by the Examiner ("to reduce ambient temperature crystallization") does not relate to block chain rearrangement. Giles teaches that short oxyalkane sequences, i.e., low values of m, are desirable to reduce ambient temperature crystallization.

Docket No.: 20241/0202402-US0

⁹ Amendment filed on May 30, 2007, page 9, last paragraph.

¹⁰ Non-Final Office Action dated March 8, 2007, page 4, lines 1-3.

¹¹ Non-Final Office Action dated March 8, 2007, page 5, lines 3-8.

Application No. 10/523,085 Amendment dated September 12, 2008 Reply to Non-Final Office Action of May 13, 2008

Docket No.: 20241/0202402-US0

Preferably, the value should lie between 7 and 17.¹² Based on this teaching, a skilled artisan that wished to reduce ambient temperature crystallization would use short oxyalkane sequences. Therefore, a skilled artisan that wished to reduce ambient temperature crystallization in the Khan ABA block copolymer, namely,

would shorten the oxyethylene side chains. Contrary to the Examiner's contention, this would not result in a rearrangement of the block chain sequence; it would only affect the length of the side chains in the **A** blocks.

At least for this reason, Applicant respectfully submits that one of ordinary skill in the art would not have been motivated at the time of the present invention to rearrange the Khan ABA block copolymer.

In addition, while the Examiner states that the "instantly claimed block chain A also could contain low values of m beginning with number 2," 13 the block chain A recited in claims 1 and 16 as amended no longer contains low values of m. (The remaining pending claims 3-6, 9, 11, 12, 14, and 15 depend from claim 1, and claims 17-20, 23-25, and 27 from claim 16.) Also, one of ordinary skill in the art would know that having long oxyalkane sequences would make it difficult for the living anion polymerization process during synthesis to proceed smoothly.

¹² Giles, column 5, lines 8-15.

¹³ Non-Final Office Action dated May 13, 2008, page 3, lines 17-18.

For this additional reason, Applicant respectfully submits that one of ordinary skill in the art would not have been motivated to rearrange the Khan block copolymers to arrive at the present invention.

Further, while the Examiner states that Giles discloses ABA triblock copolymers, Giles neither discloses nor teaches examples demonstrating that ABA triblock copolymers with a microphase separated structure can be prepared actually. Giles discloses the so-called "iniferter" method as a method to be adopted for synthesizing such triblock copolymers.

14 The term "iniferter" is an abbreviation of "initiator-transfer agent-terminator, "15 and the iniferter method is one of living radical polymerization methods.

Although the iniferter method allows the radical polymerization to take place in a living manner, the control of the polymerization is quite difficult. Therefore, the iniferter method tends to broaden the molecular weight distribution of the resulting polymers and cause termination of the polymerization or dimerization. For this purpose, Applicant submits an article taken from pages 121-126 of the book titled "Radical Polymerization Handbook" as Exhibit A, and an English translation of the article as Exhibit B. The article was authored by T. Otsu. In introducing the iniferter method, Giles cites a journal article also co-authored by T. Otsu. The Radical Polymerization Handbook was first published in 1999.

In contrast to the iniferter method, it is well known to one having ordinary skill in the art that the formation of a microphase separated structure requires a narrow molecular weight distribution and a controlled molecular weight.

Accordingly, Applicant respectfully submits that it would not have been obvious at the time of the present invention for one having ordinary skill in the art to obtain a copolymer having a microphase separated structure by incorporating polymethacrylate with an oligo(oxyethylene) block

¹⁴ Giles, column 7, lines 9-11.

¹⁵ Exhibit B, page 3, lines 8-9.

¹⁶ Giles, column 7, lines 9-11.

in the middle in accordance with the methods disclosed or taught by Khan or Giles to arrive at the subject matter of claim 1 or 16. The remaining pending claims depend from claim 1 or 16.

At least for the reasons stated above, Applicant respectfully submits that Khan in view of Giles does not render any of claims 1, 3-6, 9, 11, 12, 14-20, 23-25, and 27 obvious, and respectfully requests that the rejections of these claims be withdrawn.

CONCLUSION

In view of the foregoing, it is believed that pending claims 1, 3-6, 9, 11, 12, 14-20, 23-25, and 27 are in immediate condition for allowance and it is respectfully requested that the application be reconsidered and that all pending claims be allowed and the case passed to issue.

If there are any other issues remaining which the Examiner believes could be resolved through a Supplemental Response or an Examiner's Amendment, the Examiner is respectfully requested to contact the undersigned at the telephone number indicated below.

Dated: September 12, 2008

By four le

Registration No.: Reg. No. 47,522

DARBY & DARBY P.C.

P.O. Box 770

New York, New York 10008-0770

(212) 527-7700 (212) 527-7701 (Fax)

Attorneys/Agents For Applicant

Exhibit A

ラジカル重合ハンドブックー

基礎から新展開まで -

<----- 監修者

蒲池 幹治

(福井工業大学工学部教授・大阪大学名誉教授)

遠藤 岡

(東京工業大学資源化学研究所所長・教授)

⟨□───編集幹事 相田 卓三 | 小林 四郎 (五十音順) 市原 祥次 | 佐藤 恒之

岡本 住男 | 澤本 光男

岡谷 卓司 山田文一郎

第4節 イニファータ法によるリビングラジカル重合

1. はじめに

1955年、Srwar"はオファレンナトリウムによる スチレンの重合からリビングボリマーが地成するこ とを報復した。このリビング減合の発見は、高分子 の一次構造(分子量、分子量分布、米痛影構造な ど)の前即やテレケリック・プロック・クファトは リマーの設計や成に大きなインケトを与えられ、 1970年頃からアニオン重合、カテオン重合、ラジ カル重合、開設合、形成の高く、メケセン、重合、 直縮合などのリビング化が造成され、さらにグルー 7移動量合、イモータル最合、解集港移向配送室 合などのリビング派が構造された。18

このうち、ラジカル集合以外のリピング系では、 新規触媒系の開発によって達成されたが、 ラジカル 重合は、電気的に中性で短寿命のフリーラジカルを 中間体(成長活体験)とする連鎖反応で進むため、 涌常の触継 (開始期) によるリピング化は期待でき ない。また、フリーラジカルには、DPPHやフェル ダジルなどのように長寿命のラジカルも存在するが. これらは重合の禁止剤として機能し、安定ラジカル と呼ばれて、ここでいうリピングラジカルではない。 リピングラジカル重合化とは、短寿命のラジカルを いかにして長寿命化するかということになる。それ には、成長ラジカルの失活(修正)を防ぐため、そ の鳥動度をゼロにすることが考えられ、乳化系、沈 殿系、凍結系、ミクロゲル系、高粘度媒体中などで の重合が試みられたが、ESR で成長ラジカルは検 出されるものの、愛合はリビング的に進行しなかっ た21-81。

このように考えてくると、均一系リビングラジカル重合を構築するためには、余く新しい経効の導入 が必要であった。1982年、大歩らはラジカル重合 で生成するボリマーの末端構造を制御し、設計する ためのイニファータ法。たらびに均一系リビングラ ジカル重合モデル³⁰を模束した。今日では、イニフ エータやそれを用いたリビングラジカル壁合という 用語は広く用いられているが、 創解されていること も多い。本情では、これらの基本的な考え力を整理 するとともに、 最近の研究との関係についても述べ る。 なお、詳細は総説⁸⁻⁴¹などを参照していただき たい。 また、ブロック・グラフトポリマーの設計会 成については別報⁶⁰でも基本であ

周知のごとく、ラジカル素会は多数のビニル系モノマーの高分子化に有効で、火量の汎用ビール系ボリマーがご業生産されている。しかし、分子量分布や一次標準(とくに立体構造)の制節はあまり進んでいない。近年来、精密素合の気速の高まりから、リビングラジカル系合などによるラジカル素合の接触がである。バルスレーザー重合による皮受ラジカルの精密構造、速度解析などが動しく展開されている。ラジカル最合のメリットを虚かした精密素合法としての過度が持たれている。

2. イニファータとは

ラジカル集合の停止は、再結合と不均化で起こり、連鎖移動が低こらない限り、それぞれ取消場と片末 環に開始期別片が結合したポリマーが生成する。 関反応の起こりやすざは主生してモノマーの構造で決 められているため、ポリマーの未満基構造を制御す ることは難しい。しかし、関始期への運動移動とー メラジカル停止のいずれか、または例すを起こしや すい関始制 (R-R) を用いた場合には、尚末端に 関始剤切片(E) が結合したポリマーが生成することになる。

R-R+#M → R-(M) R

この反応は、開始剂へのモノマーの挿入反応には かならない。このような操能を持つ開始刺に対して イニファーケ (initiator-transfer agent-terninator の助) と名づけられた。そのルーツは、Gomberg によるトリテルラジカル (1) かその二量体 (2) と の解離平衡にあること、これにビニルモノマーが挿 入される (3) という Marvel らの研究であろう。 式2は4、外電で流へるリビンテラジカル像やモアル

ita i j.
i).
Chem.

iles, 30,

Irs. 28,

er and

会予精 得本

1d K.

zek i

\ 5643

ierg :

: 高分 v. 31.

d M.

nto :

ny : Wei:

Nei:

J. W.

₹>

Cl_C-CI+#M -- Cl_C -(M)-- Cl

1982年、イニファータ法を提案したときに用い たイニファータは、フェニルアゾトリフェニルメタ ン(4)、テトラフェニルエタン類(X=CgH₃ CN、 OCsH₃ など)(5)、テトラエチルチウラムジスルフ ィド(6) のようなイオウ化合物である⁸⁷。 前二者 は熱、後者はポイニファータとして機能する。

$$(C_2H_1)_2NCS - SCN(C_2H_1)_2+nM -$$
 \parallel
 S
 S

$$(C_2H_2)_3NCS \longrightarrow (M)_2 \longrightarrow SCN(C_2H_3)_2$$

 \parallel
 \parallel
 \parallel
 \parallel
 \parallel

ここで、 生成したポリマー (4-6) の水積輸は、 ち らに然あるいは光イニファータ (高分チイニファータ (高分チイニファータ (高分チイニファータ) として働くことが見いだされた。この場合のイニファータ結合は、 それぞれCーC(トリチル)、 CーC(ジフェニルメチル)、 CーS結合である。 競近、 往目をれているスチャレン/ニトロキンドリビング系では、 C(には) - ON 場合である¹⁸。 反応はこれにモノマーの婦人反比で選む。このような高分チイニファータにより第二エノマーの異をを行うことによって種々のプロックポリマーが実成することになる。 粉ぱ (5) から 110 74% 61 75。

$$9+mM'\frac{h\nu}{r} \frac{C_0H_0}{C_0H_0} \stackrel{N-C-S-(M)_0-(M')_0}{\sim} M' \stackrel{N-C-N}{\sim} \frac{C_0H_0}{r} \stackrel{(7)}{\sim} (C_0H_0)$$

このように、種々のイニファータを用いることに より官能性、テレケリック、プロック、グラフトポ リマーなどが設計合成されることになる (4.項参 照)。

8. 均一系リビングラジカル重合モデル

式22、(4)-(7)に示したように、イニファータ結合 にモノマーが挿入して進む場合には、このラジカル 最合はリビング的に進行することになる。モノマー の挿入は、イニファータ結合の解集。モノマーの付 加・ルテシフルが停止(PRT)で起こる。1982年、 提案したイニファータを用いて均一項リビングラジ カル東合モデルを式(8)に示す。ここで、CT はイニ ファータ結合(C一B)への迷鏡が移動で、この反応 が起こっても式8)のモデルには関係しない。

ここで、10のピーB結合の解離により11と18が生成する。11は成長ラジカルでMと反応し、12は原則として開始能を持たないが、11とのみ反応 PRT) し10に図る。分かりやすく書くと、次9となる。

したがって、10のC-B結合は11の休止 (dormant) 権と見られる高分子イニファータである。このモデルに従って重合が超こるなら分子量は 時間(重合率)とともに増大(図1)し、生波ポリ 義合権区

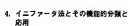
図1 2 合

⊠2 21



とになる。このため、ラジカル反応で構造制御を考えるとき、まず11の選択性を高めるために低い温度で反応を行うのが原則である。4 やちのような熱イーファータでは、70~100℃ で重合を誘きなめ、11 と B・中のペンゼン核への復奏が副反応として起こり、リビング性は低下する。6 のようなイオウ系光イニファータ (~C-S) では、実温で重合が進行するため副反応は差けられるものの、近9の11 と M の反応(成長)に比して11 と B・との反応(PRT)が遅いために、成長が連携的に進み、単分数に近いポリマーは得られない。

1994年、Georges ら¹¹¹は、2.2.6.6・ナトメメル ピペリジニル・1-オキシル(TEMPO)、連酸化ペン ソイル、St を 95 ℃で3.5 加熱し、ついで 183 ℃ 69 私食合ぎせることによって M_e/M_e-1.27 のポリ St を含成した。第一段の 反応でイニファーク結合 ので(CAは)・ON、が生成し、この結合に St モノマー が構入されて遊む。この総合は高い結合解離エネル ギーを持つこはいえ、123 ℃、86 hの 灰底は会く子 想を超えるものであったが、構造的に 製成砂が関密 されており、かつ B・は安定ラジカル(禁止制)で あるため、上記リビングラジカル風合モデルの条中 が構たさま、関・ジカル風合モデルの条中 で構たさると関で影響にしたと考えられ る。現代、多くの研究的によって研究されており、 新たなると関係と繋げらない。



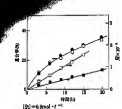
イニファータを用いて高分子の末端無構造の設計 ならびにプロックポリマーなどを設計合成する方法 はイニファータ法と呼ばれる。ここでは、われわれ が研究してきたイオウ系光イニファータ法を中心に 述べる^{2・1}。

イニファータは機能的に二つのタイプ、A-B型 とB-B型に分類される。

A-B = A+ + B - M - A - (M + B

ここで、A・は高反応性のラジカル、B・は非(または低) 反応性ラジカルで、それぞれ開始と停止に 寄与する。

B-B = B++B = M B - (M)- B



(21]=7.8×10⁻³mol・ℓ⁻¹(●、■) (22]=3.8×10⁻³mol・ℓ⁻¹(○、□)、20℃ 1 21またが22ま田いた9が光重会における時

図1 21および22を用いたStの光重合における時間-重 合率および時間-分子量の関係

マーはブロックポリマーなどの合成に用いられる。 4~6をイニファータとしたスチレン (St) ヤメタク リル酸メチル (MMA)のリピングラジカル電合がこ の例である。

これら残合を、リビングイオン賞合と比較すると、 生成ポリマーの分子装の増大、プロックポリマーの 地域では一般するが、分子整分者は単分数に近づか ない (図2参照)。イオン系合では開始が成長に比 して送やかに起こり、二分子符止も起ことないのに 対し、ラジカル家合では開始が返く、成長は達く、 10の機度を高めると二分子符化も等為に起こるこ

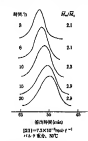


図2 21による光重合で得られたポリステレン(St)の GPC 推練

の 休 止 - タであ み子量は k 成ポリ

(9)

もことに ラフトポ

(4.項卷

:デル

- 夕結合

ラジカル

モノマー

マーの付

1982年.

・グラジ `はイニ

:の反応

ET/CT

H₂=CHX TRT/CT

11 년 12

St., 12

つみ反応

パ(9)と

(8)

第2編 ラジカル乗会屋場一屋店設計と精密業会

ここで、B-は低反応性ラジカルで開始と停止の 汲方に関与する。いま、B-B型のジスルフィド6 を光イニファータとして St を重合した場合には、 13 のポリマーが生成する。

13の両末端には、イニファータ落が結合しているが、一つ手前のモノマー単位まで考えると末端基 構造 14,15は同じではなく、光解離の結合位便も異なる。すなわち、

右(h) 來端卷:

左(1) 家羅基:

矢印の総合で解離すると、14 末編からは成長ラジカル 11 であるポリスチリルラジカル 16 を将生し、 関助に低反応のチイルラジカル 17 を生成する。し かし、15 末端からは低反応性のポリフェニルエチ ルチィルラジカル 18 と高反応性の決案ラジカル 19 が生成し、11 は再生されない。これでは、末選際 遊設計には返当ではなく、必ず 11 が再生する A-B 型イニファータを使用しなければならない。一般に、 A-B 型の方がリビング性およびプロック効率は高

60

mill b

281 3

1997).

3 P. C.

表1に、イニファータの機能、応用値かちの分類 を示す。

イニファータによるリビングラジカル重合の特徴

5.1 熱イニファータ4と5による量合

1)	熱イニファータ(熱的に機能するもの)	4, 5 c C
2)	光イニファータ(光化学的に機能するもの)	5, 6, 21, 22 など
3 J	高分子イニファータ (高分子に含まれる結合が熱イニファータとして 機能するもの)	5、4 から得られた MMA のポリマーなど (例えば 8 など)
4)	高分子光イニファータ (高分子に含まれる結合が光イニファータとし で機能するもの)	6, 21, 22 から得られたポリマーなど (例えば 9, 10, 13 など)
5)	一官能性イニファータ (高分子の片束瘤に機能基を導入するもの)	21 22
6)	一官能像 高分子イニファータ(AB 型ブロックポリマーの合成用)	21 から得られた St のポリマーなど
7)	二宮能性イニファータ (高分子の両末端に機能基を導入するもの)	22 など
8)	二官能性高分子イニファータ (ABA 型ブロックポリマーの合成用)	22 から得られた St のポリマーなど
9)	三一多官能性イニファータ (スター、構型ポリマー合成用)	23, 25 & &
10)	多官能性高分子イニファータ(高分子中に多くのイニファーダ器を持	
	つもので、スターあるいはグラフト、架橋ポリマーの合成用)	24 & ど
11)	モノマーイニファータ(集合性の二重結合と光イニファータ能を持つ	
12)	もので、マクロマー合成やそれを用いたグラフト共産合体合成用) 高分子ケルイニファーク (異なるプロック単位を多く特つプロック共	25 ♦ €
	黄合体の合成目)	26 A P

第2条 リピングラジカル乗介による精密部割

ポリ MMA 水端:Z=CH₃、Y=COOH₅、X=CN or C₆H₅ ボリSt 水路: Z=H. Y=Cells, X=CN or Cells

ポリマーを与える。しかし、リビング性。プロック 効率はあまり高くない。

5.2 ジチオカルパメート系光イニファータに よる電合

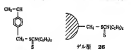
4. 項で述べたように、6はB-B型の光イニフ ァータであり、6とそれからの生成ポリマー(高分 子光イニファータ) 9とはイニファータ結合が異な る。これは、高分子設計上好ましくない。加えて、 開始反応からくる複雑さを避けるためには、成長末 端と同じイニファータ結合のものを用いればよい。 次に示すような種々の官能性 A-B 型ジエチル、ジ

$$(C_2H_3)_n(C_5-CH_2) = CH_2 - SCN(C_4H_3)_c$$

$$(C_2H_3)_n(C_5-CH_3) = CH_2 - SCN(C_4H_3)_c$$

$$(C_2H_3)_n(C_5-CH_3) = CH_2 - SCN(C_4H_3)_c$$

四官能性(架橋性) 28



重合性

チオカルパメート系光イニファータが合成され、用 いられた(機能と応用は表1齢順)。

21-26 はいずれも光イニファータとしてリビン グラジカル重合を誘起した。例えば、イニファータ 結合当たり同じ濃度の21と22を用いてStの電合 を行った際の時間一致合率関係を図1に示す。これ から、両者は同じ曲線上にある。分子量はいずれも 時間とともに増大するが、二官能性の22では、一 官能性の21の2倍となる。何じように四官能性の 23 ではゲル化の起こることが認められた。ただし、 図2に示すように生成ポリマーの分散度は通常のラ ジカル重合と回程度 (M_/M。=2) であるが、21 の 場合にはいくらか増大する傾向にあった。また、式 (8)、(9)で 11 と 12 の反応を促進するため 6 の添加を 試みたが、リピング性はいくらか向上するにとどま った。ジチオカルバメート系光イニファータでの B・はチイルラジカルであり、低いながら開始能を 持つためと考えられる。この点が、スチレン/ニト ロキシド系と異なる。

6. 星型。 ブロック、グラフト。 架橋 ポリマーの設計合成

表1に示したように、種々の官能性ジチオカルバ メート系光イニファータを用いることによって上紀 の各種ポリマーを設計合成することができたが、分 子量分布の制御はできなかった。このうち、ブロッ ク. グラフトボリマーの合成については別稿¹³で述 べる。

7. おわりに

これまで種々のイニファータ、とくにジチオカル バメート系光イニファータを用いた窓分子合成につ いて述べた。

その結果、各種ポリマーの設計合成13が可能とな った。最近の成替12)では、一つの章として取り上

125

カル19 灾蜡楼 ã A-B

一般に. 寒は高

の分類

ラジカ

型の熱 4MAの TERM あるSt 僧地エ 行する。 1) がた ル解離 、磁かめ (1) #3 (1)

いした 维性高 ロッケ

-22 مغج

2 種 ラジカル集合反応−反応設計と精密重合

げられ、また医用・機器材料へのイニファーを法の 応用も進んでいる[™]。

しかし、現在のジチオカルバメート系光イニファータ法では緩密に式場のリビングラジル集合モデルどもりに接触さず、一部側反応を伴い、結外をして分子量分布の規制は不可能であった。これでは精密流をとはいいがたい。近年来のステレン/ニトロキシド系条イニファータの使用は sk の 利密合成まで可能となった。これは、また他のリビングインクラジカル重合から単分散に近いボリ sk の利密合成まで可能となった。これは、また他のリビングイン主合の大健食このイニファータ系に変換し、リビングラジカル重合することによって分子量分布の制御まれた減分子の特密合成への直を開いた。今後の一般の経歴を開けたた。

【参考・引用文献】

- 1) M. Szwarc : J. Am. Chem. Soc., 78, 2656 (1956).
- 2) 大津陸行:高分子(総裁), 37,248 (1988).
- 大洋隆行、松本章·;高分子(轮流), 41,858 (1992).
- 4) 大津險行,松本章…, 古岡正裕:化学裁提(総説),

- 18. 精密集合、pp.3~18 (1993),
- 5) T. Otsu and A. Matsumoto: Macromol, Design (総数): Concepts and Practice, Polym. Front, Int't. Inc pp. 471-486 (1994): Adv. Polym. Sci. (総数), 136, 75-127 (1998).
- 6) 松本章--:高分子加工 (総説), 46, 338-346 (1997).
- 7) 例えば ACS Polym, Prep., 38 (1, 2) (1997) および高分子予格数, 46 (1997).
- 8) 梶凉篤, 顔池幹治: 高分子, 47,62 (1998).
- T. Otsu and M. Yoshida : Makromol . Chem., Rapid Commun., 3, 127-132 (1982).
- T. Otsu, M. Yoshida and T. Tazuki : ibid., 3, 133-140 (1982).
- M. K. Georges et al.: Trends Polym. Sci., 2, 66 (1994);
 Macromol. Symp., 88, 89 (1994).
- Misra ed. : Macroniol Design : Concepts & Practice, Polymer Frontier Int'l. Inc. (1994).
- 13) 本書第2編第3章第2節2.項,

<大津 隆行>

11 12:

り開発機

配位为数

神の変配を 中華分類が 海線に対す たけビング

ラジカル筆合ハンドブックー基礎から新展開までー

定価 56,400 円 (統別) ©1999 (統印省略)

綱 終 松風まさみ 製 作 略オフィス東和 印 駒 金昭和登印彫所

発行者 盲田 陰 :SBN4-900839-41-0

企画・標準 整発エヌ・ティー・エス

iSBN4-000830-41-0 〒113-8755 東京移文京区播海 2-16-16 は203(8814)5511間の ※低丁・茶丁本はお取り替えします。鉄筋観写・転載を繋す。 企画・柳原 高麗エヌ・ティー・エス 姓 行 芝菜エヌ・ティー・エス

Exhibit B

LIVING RADICAL POLYMERIZATION USING INTERTER METHOD

1. Introduction

In 1956, Szware¹⁾ reported that a living polymer is generated from the polymerization of styrene using sodium naphthalene. The delivery of the living polymerization had a great impact on the control of the primary structure (such as molecular weight, molecular weight distribution, and end group structure) of macromolecules or the design and synthesis of telechelic, block, and graft polymers. Thereafter, the living polymerization was considered as the origin of the structure control, the living of anion polymerization, cation polymerization, radical polymerization, ring-opening polymerization, coordination polymerization, metathesis polymerization, and polycondensation was accomplished from 1970, and the living systems such as group transfer polymerization, immortal polymerization, and new transition metal catalyst polymerization were reported²²⁻⁵⁾.

The living systems other than the radical polymerization were constructed with the development of a new catalyst system. However, the living radical polymerization could not proceed in a living manner in the presence of a usual catalyst (initiator), because the radical polymerization proceeded via chain reaction while forming electrically-neutral short-lived free radical intermediates as growth-activated radicals. Although there are long-lived free radicals such as DPPH or verdazyl, the radicals serve as polymerization inhibitors and are called stable radicals, which are not the living radicals mentioned herein. The living radical polymerization relates to how to elongate the lifetime of a short-lived radical into a long lifetime. In order to prevent the deactivation of the growth radical by making the mobility of the living radical zero, the polymerization was tried in an emulsion system, a precipitation system, a freezing system, a micro gel system, or a high-viscosity

medium. However, the growth radicals were detected in ESR but the living polymerization did not proceed in a living manner²⁾⁻⁸⁾.

From this point of view, the introduction of a new idea was required to construct the homogeneous-system living radical polymerization. In 1982, OTSU et al. suggested an iniferter method⁹⁾ and a homogeneous-system living radical polymerization model¹⁰⁾ for controlling and designing the end group structure of polymers generated by the radical polymerization. In recent years, the terms "iniferter" or the "living radical polymerization" using the iniferter were widely used but often misunderstood. In this description, the basic way of thinking thereabout is arranged and the relation thereof to the recent studies is also described. The details thereof is referred to in the Introduction²⁾⁻⁸⁾. The design and synthesis of the block and graft polymers will be described in another document¹⁹⁾.

As widely known, the radical polymerization is effective for the polymerization of vinyl monomers and a large amount of general-purpose vinyl polymers are industrially produced. However, the control of the molecular weight distribution or the primary structure (particularly, stereoscopic structure) was not satisfactorily advanced. In recent years, the structure analysis of the radical polymerization such as the living radical polymerization or the precise structure or speed analysis of the growth radicals in the ESR / pulse laser polymerization have been actively developed. Accordingly, the advancement as the precise polymerization method having the merit of radical polymerization is expected.

2. Iniferter

The radical polymerization is stopped due to the recombination and nonhomogeneousness, and polymers in which initiator pieces are coupled to both ends or one end thereof are produced unless the chain transfer occurs. Since the occurrence of both reactions depends on the monomer structure, it is difficult to control the end group structure of the polymers. If an initiator (R-R) easily causing at least one of the chain transfer to the initiator and the primary radical stop is used, polymers in which pieces of the initiator (R) are coupled to both ends are formed.

This reaction is only an insertion reaction of monomers into the initiator. The initiator having such a function was named an "iniferter" (abbreviation of <u>initiator-transfer</u> agent-<u>ter</u>minator). Formula 2 illustrates the living radical polymerization model described below. The function of telogen (chain transfer agent) in telomerization results from the insertion reaction of monomers. For example, when CCl₄ is used as telogen, the reaction shown by Formula (3) occurs.

ChC - C1+#M -- ChC -- (M) -- CI

An iniferter used in 1982 was a sulfur compound such as phenyl azotriphenyl methane (4), tetraphenyl ethane (X=C₂H₅, CN, OC₆H₅, or the like) (5), or tetraethyl thiuram disulfide $(6)^{8/7}$. The former two compounds serve as a thermal iniferter and the latter serves as an optical iniferter.

Here, it was also discovered that the end groups of the produced polymers (4) to (6) serve as the thermal or optical iniferter (referred to as macromolecule iniferter). The iniferter bond in each case is C-C (trityl), C-C (diphenyl methyl), or C-S bond. In recent years, the iniferter bond in the living system of styrene/nitroxide is C(C₆H₅)-ON- bond¹¹. The reaction is the insertion reaction of monomer thereto. By polymerizing a second monomer using the polymer iniferter, various block polymers are produced. For example, 10 are obtained from 9.

In this way, functional, telechelic, block, or graft polymers are designed and synthesized by using various iniferters.

Homogeneous Living Radical Polymerization Model
 As shown in Formulas 2 and 4 to 7, when monomers are incorporated in the

iniferter bonds in radical polymerization, the radical polymerization proceeds in a living manner. The insertion of monomers occurs with the dissociation of the iniferter bonds, the addition of monomers, or the primary radical termination (PRT). The homogeneous living radical polymerization model using the iniferter suggested in 1982 is expressed by Formula 8. Here, CT is a chain transfer to the iniferter bond (C-B) and the reaction is not associated with the model expressed by Formula 8.

Here, 11 and 12 are produced by the dissociation of C-B bond of 10. 11 is a growth radical which reacts with M. 12 does not have an initiation function in principle, but reacts with only 11 to return to 10. as shown by Formula 9.

Thus, the C-B bond of 10 is a polymer iniferter deemed as a dormant seed of 11.

When polymerization occurs in this model, the molecular weight increases with time
(polymerization rate) (FIG. 1), and the produced polymer may be used to synthesize block
polymers or the like. An example thereof is the living radical polymerization of styrene
(St) or methyl methacrylate (MMA) using 4 to 6 as iniferter.

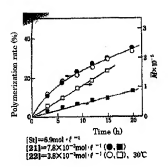
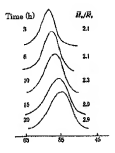


FIG. 1 Relations between the time-polymerization rate and between time and molecular weight in photopolymerization of St using 21 and 22

Although these polymerizations are coincident with living ion polymerization in terms of the increase in molecular weight of the produced polymers and the production of the block polymers, the molecular weight distribution is not close to mono-dispersion (see FIG 2).



Elution Time (min)
[21] = 7.3 × 10⁻³ mol/l
Bulk polymerization at 30°C

FIG. 2 GPC curve of polystyrene (St) obtained by photopolymerization using 21

In the ion polymerization, the initiation reaction occurs rapidly in comparison with the growth reaction and no bimolecular termination occurs. In contrast, the initiation reaction is slow, the growth reaction is rapid, and the bimolecular termination also easily occurs in accordance with the increase in concentration of 11 in the radical polymerization.

Accordingly, the structure control in the radical reaction requires that the reaction is carried out at a low temperature to enhance the selectivity of 11. Since the thermal iniferters such as 4 or 5 triggers the polymerization at 70°C to 100°C, the substitution of 11 and B in benzene nucleus occurs as the side reaction and the living property is deteriorated. Sine the polymerization using the sulfur-based photo-iniferter (—C-S) such as 6 proceeds at a room temperature, the side reaction can be avoided. However, since the reaction of 11 and

B · (PRT) proceeds slowly in comparison with the (growth) reaction of 11 and M of Formula 9, the growth reaction proceeds in chains and the polymers close to the monodispersion cannot be obtained.

In 1994, Georges et al. ¹¹ synthesized poly St with M_w/M_a =12.7 by heating 2,2,6,6,tetramethylpeperidinyl-1-oxyl (TEMPO), benzoil peroxide, and St at 95°C for 3.5 hours
and polymerizing them at 123°C for 69 hours. The iniferter bond ($C(C_6H_3)$ -ON-) is
produced in the first stage of the reaction and a St monomer is inserted into the bond.
Although the bond has high bonding-dissociation energy, the reaction at 123°C for 69 hours
exceeds the expectation, the side reaction is hindered in structure, and B • is a stable
radical (inhibitor). Accordingly, it is considered that the condition for the living radical
polymerization is satisfied and thus the production of the polymer (here, poly St) exhibiting
approximate mono-dispersion is realized by the radical polymerization. A lot of
researchers are currently dedicated to the study and new developments are expected 6,8).

4. Iniferter Method and Functional Classification and Application thereof

The method for designing an end group structure of a polymer and designing a block polymer using an iniferters is called the iniferter methods. A sulfur-based photoiniferter method will be described below.²⁾⁻¹⁾

Here, $A \cdot is$ a high-reactive radical and $B \cdot is$ a non- (or low-) reactive radical, where contribute to the initiation and termination, respectively.

Here, B • is a low-reactive radical and contributes both the initiation and termination reactions. When St is polymerized using disulfide 6 of the B-B type as the

photo-iniferter, polymer 13 is produced.

Iniferter groups are coupled to both ends of 13. However, the end group structures 14 and 15 are different from each other in terms of the previous monomer unit, and the bonding positions to be photodissociated. That is,

Right (h) end group:

Left (t) end group;

If a bonding of 14 shown by the arrow is dissociated, polystyryl radical 16 as the growth radical 11 is reproduced from the end of 14 and low-reactive thiyl radical 17 is produced at the same time. Low-reactive polyphenylethyl thiyl radical 18 and high-reactive carbon radical 19 are produced from the end of 15 and 11 is not reproduced. Accordingly, this is not suitable to design the end structure, and the iniferter of A-B type by which 11 is necessarily reproduced should be used. In general, the living property and the block efficiency of the A-B type are higher.

The classification of iniferters in view of functionality and application is shown in Table 1.

Table 1 Classification and Application of Iniferters in view of Functionality

1	Thermal Iniferter (thermally functioning)	4, 5, or the like.
2	Photo-iniferter (photochemically functioning)	5, 6, 21, 22, or the like.
3	Polymer iniferter (bond of polymer functions as a thermal iniferter)	MMA polymers (such as 8) obtained from 5 or 4, or the like.
4	Polymer iniferter (bond of polymer functions as a phot- iniferter)	Polymers (such as 9, 10, or 13) obtained from 6, 21, or 22, or the like.
5	Mono-functional iniferter (functional group is introduced into one end of a polymer)	21, or the like.
6	Mono-functional polymer iniferter (for synthesis of AB type block polymer)	St polymers obtained from 21, or the like.
7	Bi-functional iniferter (functional groups are introduced into both ends of a polymer)	22, or the like.
8	Bi-functional polymer iniferter (for synthesis of ABA type block polymer)	St polymers obtained from 22, or the like.
9	3 or more functional iniferter (for synthesis of star or comb type polymer)	23, 25, or the like.
10	Multi-functional polymer iniferter (having many iniferter groups in a polymer and available for synthesis of star, graft, or cross-linking polymers)	24, or the like.
1	macromer or synthesis of a graft copolymer using it)	25, or the like
12	Polymer gel iniferter (for synthesis of a block copolymer having many different block units)	26, or the like.

- 5. Features of a Living Radical Polymerization using an Iniferter
- 5.1 Polymerization using Thermal Iniferters 4 and 5 Thermal iniferters 4 and 5 are A-B type and B-B type of thermal iniferters,

respectively, and trigger the radical polymerization of St or MMA at 70°C to 100°C (Formulas 4 and 5), and produce polymers 7 and 8. The polymerization of St which is a mono-substituted ethylene does not proceed in the living manner, but the polymerization of MMA which is a 1,1-di-substituted ethylene proceeds in the living manner. The study using the model compounds demonstrated that the radical dissociation of the right (h) end group bond (20) of 7 or 8 having a 6-substituted ethane structure occurred in MMA. The radical dissociation of St having a 5-substituted ethane structure, or a left (t) poly St or MMA ends having a 4-substituted ethane structure did not occur at 70°C to 100°C.

Accordingly, the polymers 7 and 8 obtained from MMA serve as mono-functional polymer iniferters and give the A-B type block polymers. However, the living property and the block efficiency are not high.

Poly MMA end: Z=CH₃, Y=COOH, X=CN or C₆H₅
Poly St end: Z=H, Y=C₆H₅, X=CN or C₆H₅

5.2 Polymerization using Dithiocarbamate Photo-Iniferter

As described above, 6 is a B-B type photo-iniferter, and 6 and polymer (polymer photo-iniferter) 9 produced therefrom are different in iniferter bond. This is not desirable for polymer design. In addition, in order to avoid the complication resulting from the initiation reaction, the same iniferter bond as that of the growth end can be used. Various functional A-B type diethyl and dithiocarbamate-based photo-iniferter described below were synthesized and used (see Table 1 for the functions and applications).

O- CHF - RENCCHOS

Mono-functional 21

$$(C_0H_0)_*N_{\mathfrak{S}}^{\mathsf{CS}} - CH_{\mathfrak{s}} - \bigcirc - \mathsf{CH}_{\mathfrak{s}} - \sup_{\mathfrak{S}} N(C_{\mathfrak{s}}H_0)_{\mathfrak{s}}$$

Bi-functional 22

$$(C_2H_2)_1NC^2S - CH_2 \qquad CH_2 - \frac{1}{8}CN(C_2H_3)_2$$

$$(C_2H_2)_1NC^2S - CH_2 \qquad CH_2 - \frac{1}{8}CN(C_2H_3)_2$$

Tetra-functional (cross-linking) 23

Multi-functional (cross-linking) 24

Polymerized 25



Gelated 26

21 to 26 are all photo-iniferters and triggered the living radical polymerization. For example, the time-polymerization relation is shown in FIG. 1 where the polymerization of St was carried out using 21 and 22 having the same concentration per iniferter bond. Both can be seen in the same curve. The molecular weight increases with time, but the molecular weight of 22 having bi-functionality is double that of 21 having monofunctionality. Similarly, it was observed that the gelation occurs in 23 having tetrafunctionality. However, as shown in FIG. 2, the degree of dispersion of the produced polymer is similar to that of the usual radical polymerization $(M_w/M_n=2)$, but tends to increase to only some extent in 21. The addition of 6 was tried to promote the reaction of 11 and 12 in Formulas 8 and 9, but the living property was improved to some extent. This is probably because B \cdot in the dithiocarbamate-based photo-iniferter is thiyl radical and has a low initiation function. In this point, the dithiocarbamate-based photo-iniferter differs from a styrene/nitroxide iniferter.

6. Design and Synthesis of Star, Block, Graft, and Cross-linking Polymers

As shown in Table 1, various polymers described above could be designed and synthesized by using various functional dithiocarbamate photo-iniferters, but the molecular weight distributions thereof could not be controlled. The synthesis of block and graft polymers would be described in another document¹³.

7. Conclusion

The synthesis of polymers using various iniferters, particularly, a dithiocarbamatebased photo-iniferter, has been described above.

As a result, the design and synthesis¹³⁾ of various polymers were possible. The iniferter method was treated as one chapter in recent written reports 12) and the application of an iniferter method to medical instrument materials has advanced ¹³⁾.

However, strictly speaking, the living radical polymerization model of Formula 8 does not function in the present dithiocarbamate-based photo-iniferter method and the control of the molecular weight distribution was not possible as the result of some side reactions. This cannot be said as precise polymerization. The recent use of styrene/nitroxide thermal iniferter enabled the living radical polymerization of St and the precise synthesis of poly St exhibiting an approximate mono-dispersion. It opens the road to the precise synthesis of polymers of which the molecular weight distributions are controlled by converting the radicals of other living ion polymerization into the iniferter groups to polymerize the living radicals. Further development is expected in the future.

[Cited Documents for Reference]

- 1) M. Szwarc: J. Am. Chem. Soc., 78, 2656 (1956)
- 2) Takayuki OTSU: Macromolecule (Introduction), 37, 248 (1988)
- Takayuki OTSU, Akikazu MATSUMOTO: Macromolecule (Introduction), 41, 358 (1992)
- Takayuki OTSU, Akikazu MATSUMOTO, Masahiro YOSHIOKA: Introduction of Chemistry (Introduction), 18. Precise Polymerization, pp.3-18 (1993)
- T. Otsu and A. Matsumoto: Macromol. Design (Introduction); Concepts and Practice, Polym. Front. Int'l. Inc. pp.471~486 (1994); Adv. Polym. Sci. (Introduction), 136,

- 75-137 (1998)
- Akikazu MATSUMOTO: Macromolecule Processing (Introduction), 46, 338-346 (1997)
- For example, ACS Polym. Prep., 38(1,2) (1997) & Preliminary Announcement of Macromolecule. 46, (1997)
 - 8) Atsushi KAJIWARA, Mikiharu KAMACHI: Macromolecule, 47, 62 (1998)
 - 9) T. Otsu and M. Yoshida: Macromol. Chem., Rapid Commun., 3. 127-132 (1982)
 - 10) T. Otsu, M. Yoshida, and T. Tazaki: ibid., 3. 133-140 (1982)
- M. K. Geoges et al.: Trends Polym. Sci., 2, 66 (1994); Macromol. Symp., 88, 89 (1994)
- 12) Misra ed.: Macromol. Design: Concepts & Practice, Polymer Frontier Int'l. Inc. (1994)
 - 13) Present Volume, Second Edition, Chapter 3, Paragraph 2, Section 2.
 <Takavuki OTSU>